```
=> FILE REG
FILE 'REGISTRY' ENTERED ON 06 MAY 2008
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2008 American Chemical Society (ACS)
=> DISPLAY HISTORY FULL L1-
    FILE 'REGISTRY' ENTERED ON 06 MAY 2008
                E CARBON/CN
L1
              1 SEA CARBON/CN
L2
        27886 SEA 50-00-0/CRN
                E UREA/CN
1.3
              1 SEA UREA/CN
               E MELAMINE/CN
L4
              1 SEA MELAMINE/CN
L5
              2 SEA L3 OR L4
                SEL L5 1-2 RN
                EDIT E1-E2 /BI /CRN
L6
         14066 SEA (108-78-1/CRN OR 57-13-6/CRN)
L7
          9301 SEA L2 AND L6
    FILE 'HCA' ENTERED ON 06 MAY 2008
          47504 SEA L7 OR (FORMALDEHYDE# OR CH20 OR H2CO OR HCH0 OR
L8
                FORMALIN#)(2A)(UREA# OR NH2CONH2 OR H2NCONH2 OR MELAMINE#
T. 9
          55654 SEA POLYISOCYANAT? OR (?ISOCYANAT? OR NCO OR RNCO)(2A)(AD
                DUCT? OR POLY OR POLYM? OR COPOLYM? OR HOMOPOLYM? OR
                RESIN?)
L10
           1094 SEA L1 (L) FOAM?
L11
           3133 SEA (CARBON# OR C OR CARBONACEOUS? OR CARBONIFEROUS?) (2A)
               FOAM?
L12
        114289 SEA PYROLY?
    FILE 'REGISTRY' ENTERED ON 06 MAY 2008
               E CARBON DIOXIDE/CN
L13
              1 SEA "CARBON DIOXIDE"/CN
    FILE 'HCA' ENTERED ON 06 MAY 2008
        517144 SEA L13 OR CO2 OR CARBON#(A)DIOXIDE#
L14
         352425 SEA STEAM? OR (WATER? OR H2O)(2A)(VAPOR? OR VAPOUR? OR
L15
               GAS## OR GASEOUS? OR GASIF?)
            88 SEA L11 AND L12
L16
L17
            8 SEA L16 AND (L14 OR L15)
L18
             1 SEA L17 AND (L8 OR L9)
```

```
L19
           43 SEA L10 AND L12
L20
             3 SEA L19 AND (L14 OR L15)
L21
             1 SEA L20 AND (L8 OR L9)
L22
        156693 SEA FOAM?
L23
           893 SEA L22 AND L12
L24
           107 SEA L23 AND (L14 OR L15)
L25
             5 SEA L24 AND (L8 OR L9)
L26
         218040 SEA POLYURETHAN## OR URETHAN## OR POLYAMIC#
L27
            42 SEA L24 AND L26
L28
          22019 SEA L13 (L) RACT/RL
L29
             0 SEA L27 AND L28
L30
             0 SEA L17 AND L26
L31
            0 SEA L20 AND L26
L32
            85 SEA L23 AND L14
L33
            29 SEA L23 AND L15
L34
             5 SEA L33 AND L26
L35
        72629 SEA (HIGH? OR RAIS? OR ELEVAT? OR INCREAS? OR GREAT?) (3A)
               (SURFACE?(2A) AREA# OR SA OR S(W) A OR PORE# OR PORO? OR
               MICROPORO? OR MICROPORE# OR M2(A)(G OR GR OR GRM# OR GM#
               OR GRAM#))
L36
          22086 SEA (CELL OR CELLS OR CELLULAR?) (2A) (SIZE# OR SIZING# OR
               DIA# OR DIAM# OR DIAMET? OR RADII? OR RADIUS?)
L37
             5 SEA L24 AND L35
L38
             5 SEA L24 AND L36
    FILE 'HCAPLUS' ENTERED ON 06 MAY 2008
T.39
           370 SEA RO!ERMUND ?/AU
L40
          2008 SEA HEMPEL ?/AU
L41
          4333 SEA HESSE 2/AU
L42
          256 SEA RUDLOFF ?/AU
L43
             1 SEA DESSEIX ?/AU
             1 SEA L39 AND L40 AND L41 AND L42 AND L43
L44
               SEL L44 1 RN
    FILE 'HCA' ENTERED ON 06 MAY 2008
L45
         153715 SEA (HIGH? OR RAIS? OR ELEVAT? OR INCREAS? OR GREAT?) (3A)
               SURFACE?
L46
         119577 SEA POROSIT?
L47
           7432 SEA OPEN?(2A)(CELL OR CELLS OR CELLULAR?)
L48
             9 SEA L24 AND (L45 OR L46 OR L47)
    FILE 'REGISTRY' ENTERED ON 06 MAY 2008
            12 SEA (101-05-3/BI OR 127-08-2/BI OR 134498-73-0/BI OR
L49
L50
             4 SEA L49 AND PMS/CI
             8 SEA L49 NOT L50
L51
             E ISOCYANIC ACID/CN
L52
            1 SEA "ISOCYANIC ACID"/CN
```

D L50 1-4 IDE

FILE 'HCA' ENTERED ON 06 MAY 2008

L53 38211 SEA L50 OR L52

L54 13 SEA L53 AND L12 AND L22

L55 2 SEA L54 AND (L14 OR L15)

L56 1 SEA L54 AND (L35 OR L36 OR L45 OR L46 OR L47) L57 24 SEA L17 OR L18 OR L20 OR L21 OR L25 OR L34 OR L37 OR L38

OR L48 OR L55 OR L56

T-58 14 SEA 1840-2002/PY, PRY, AY AND L57

=> FILE HCA

FILE 'HCA' ENTERED ON 06 MAY 2008

USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

=> D L58 1-14 BIB ABS HITSTR HITIND

L58 ANSWER 1 OF 14 HCA COPYRIGHT 2008 ACS on STN

140:274741 HCA Full-text AN

TΙ Manufacture of carbon composed foams with high internal surface by pyrolysis of N-containing polymers

Rotermund, Udo: Hempel, Renate: Hesse, Michael: Rudloff, Jan: IN Desseix, Maryline

PA BASF AG, Germany

SO Ger. Offen., 17 pp.

CODEN: GWXXBX

DT Patent

LA FAN.	German CNT 1				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE 10243240	A1	20040325	DE 2002-10243240	200209 17
	WO 2004026792	A1	20040401	< WO 2003-EP9943	200309 08

	W:	CN, GE, LC, NI, SL,	CO, GH, LK, NO,	CR, GM, LR, NZ, TJ,	CU, HR, LS, OM,	CZ, HU, LT, PG,	AU, DE, ID, LU, PH, TR,	DK, IL, LV, PL,	DM, IN, MA, PT,	DZ, IS, MD, RO,	EC, JP, MG, RU,	EE, KE, MK, SC,	ES, KG, MN, SD,	FI, KP, MW, SE,	GB, KR, MX, SG,	GD, KZ, MZ, SK,
	RW:	GH, BY, EE, SI,	GM, KG, ES,	KE, KZ, FI, TR,	MD, FR, BF,	RU, GB,	MZ, TJ, GR, CF,	TM, HU,	AT, IE,	BE, IT,	BG, LU,	CH, MC,	CY, NL,	CZ, PT,	DE, RO,	DK, SE,
AU	2003			10,	A1		2004	0408	i	AU 2	003-:	2663	63		2	00309
EP	1542	941			A1		2005	0622	1	EP 2	<	7972	84		2	00309
EP	1542: R:	AT,				DK,	2006 ES, FI,	FR,								
CN	1681	748			A		2005	1012	(CN 2	003-	8220	42		2	00309
JP	2006	5122	65		T		2006	0413		JP 2		5370	12		2	00309
AT	3468	32			T		2006	1215	i	AT 2	003-	7972	84		2	00309
ES	2277	148			Т3		2007	0701	1	ES 2	003-	7972	84		2	00309
US	2006	0014	908		A1		2006	0119	1	JS 2		5269	30		2	00503 8
PRAI DE WO	2002- 2003-			0	A W		2002		<	-	<					

```
walls referred to this cell size which contain in the cross section a
     triangle, and pores in the cell scaffolding material with dimensions
     from 0.2 nm to 50 nm and a vol. 0.01-0.8 cm3/g. The procedure for
     producing this foam is based on pyrolysis of synthetic foams, whereby
     the assigned plastics foam materials (e.g., urea- formaldehyde
     copolymer or melamine- formaldehyde copolymer) possess at least 30%
     of a polymer (e.g., polyisocyanate) with a nitrogen content over 6%
     and a porosity 35-995%, as well as an open-cell character >1%. The
     pyrolysis is carried out with steam, carbon dioxide, and/or oxygen at
     ≥400-1200°. The pyrolyzed plastics foam materials may comprise an
     inorg, component such as ag. dispersion of an inorg, salt (e.g.
     ZnCl2, CaCO3, or ammonium polyphosphate), metal powder, or graphite.
     The resulting carbon foam is suitable for filters, thermal
     insulators, carriers, or semi-product for further processing in
     electrodes, superconductors, or fuel cell materials.
ΙT
     7440-44-0. Carbon, uses
        (carbon foam: manuf. of carbon
        composed foams with high internal
        surface by pyrolysis of N-contg. polymers)
     7440-44-0 HCA
RN
CN
     Carbon (CA INDEX NAME)
С
ΙT
     9003-08-1, Melamine-formaldehyde
     copolymer 9011-05-6, Urea-formaldehyde
     copolymer
        (pyrolyzed plastic; manuf. of carbon composed
        foams with high internal surface by
        pyrolysis of N-contq. polymers)
RN
     9003-08-1 HCA
     1,3,5-Triazine-2,4,6-triamine, polymer with formaldehyde (CA INDEX
CN
     NAME)
     CM
          1
     CRN 108-78-1
```

The invention concerns a foam contg. 270% of carbon and having a middle cell size over 20 µm, a porosity 35-99.5%, as well as opencell character over 90%, an internal surface over 50 m2/g with cell

AR

CMF C3 H6 N6

CM 2

CRN 50-00-0 CMF C H2 O

H2C==0

RN 9011-05-6 HCA CN Urea, polymer with formaldehyde (CA INDEX NAME)

CM 1

CRN 57-13-6 CMF C H4 N2 O

CM 2

CRN 50-00-0 CMF C H2 O

H2C==0

```
foams with high internal surface by
       pyrolysis of N-contq. polymers)
RN
     75-13-8 HCA
CN
     Isocyanic acid (6CI, 8CI, 9CI) (CA INDEX NAME)
ΙT
    134498-73-0, Lupranate M 50 674292-37-6, Lupranate
        (pyrolyzed polymer; manuf. of carbon composed
       foams with high internal surface by
       pyrolysis of N-contq. polymers)
RN
    134498-73-0 HCA
CN
    Lupranate M 50 (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    674292-37-6 HCA
RN
    Lupranate M 70 (9CI) (CA INDEX NAME)
CN
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
IC
    ICM C08J009-22
    ICS C08J009-36; C08J011-12
    57-8 (Ceramics)
    Section cross-reference(s): 38
ST
    carbon foam ureaformaldehyde
    polyisocyanate pyrolysis
ΙT
    Polyphosphoric acids
        (ammonium salts, additive; manuf. of carbon composed
        foams with high internal surface by
       pyrolysis of N-contq. polymers)
     Electrodes
IΤ
    Filters
    Fuel cells
     Superconductors
     Thermal insulators
        (carbon foam for; manuf. of carbon
        composed feams with high internal
       surface by pyrolysis of N-contg. polymers)
TΤ
    Foams
      Porosity
        (carbon foam; manuf. of carbon
       composed foams with high internal
        surface by pyrolysis of N-contg. polymers)
ΙT
    Polysiloxanes, uses
        (di-Me, hydroxypropyl Me, ethoxylated propoxylated, Tegostab
        B8461, surfactant; manuf. of carbon composed
```

```
foams with high internal surface by
        pyrolysis of N-contq, polymers)
ΙT
    Polysiloxanes, uses
        (polyoxyalkylene-, Tegostab B8409, surfactant; manuf. of
        carbon composed foams with high
        internal surface by pyrolysis of N-contg.
        polymers)
IΤ
    Polyoxyalkylenes, uses
        (polysiloxane-, Tegostab B8409, surfactant; manuf. of
        carbon composed foams with high
        internal surface by pyrolysis of N-contg.
        polymers)
ΙT
    Thermal decomposition
        (pyrolysis; manuf. of carbon composed
        foams with high internal surface by
        pyrolysis of N-contq, polymers)
ΤТ
    Aminoplasts
        (pyrolyzed plastic; manuf. of carbon composed
        foams with high internal surface by
        pyrolysis of N-contg. polymers)
    Amines, uses
ΙT
        (triamines, propellant; manuf. of carbon composed
        foams with high internal surface by
        pyrolysis of N-contq. polymers)
     471-34-1, Calcium carbonate (CaCO3), uses 7646-85-7, Zinc chloride
IΤ
     (ZnCl2), uses 7782-42-5, Graphite, uses
        (additive; manuf. of carbon composed foams
        with high internal surface by
        pyrolysis of N-contq. polymers)
     7440-44-0, Carbon, uses
TΤ
        (carbon foam; manuf. of carbon
        composed foams with high internal
        surface by pyrolysis of N-contq. polymers)
ΙT
     127-08-2, Potassium acetate
        (catalyst; manuf. of carbon composed foams
        with high internal surface by
        pyrolysis of N-contq. polymers)
     101-05-3, Triazin
IΤ
        (propellant; manuf. of carbon composed feams
        with high internal surface by
        pyrolysis of N-contq. polymers)
     9003-08-1, Melamine-formaldehyde
IΤ
     copolymer 9011-05-6, Urea-formaldehyde
     copolymer
        (pyrolyzed plastic; manuf. of carbon composed
        foams with high internal surface by
        pyrolysis of N-contq, polymers)
```

IT 75-13-80, Isocyanic acid, esters, polymers (pyrolyzed polymer; manuf. of carbon composed foams with high internal surface by pyrolysis of N-contq. polymers)

IT 134498-73-0, Lupranate M 50 674292-37-6, Lupranate M 70

(pyrolyzed polymer; manuf. of carbon composed foams with high internal surface by pyrolysis of N-contq. polymers)

IT 215512-68-8, Ortegol 501

(stabilizer; manuf. of carbon composed foams with high internal surface by pyrolysis of N-contg. polymers)

- L58 ANSWER 2 OF 14 HCA COPYRIGHT 2008 ACS on STN
- AN 138:90783 HCA Full-text
- TI Polyurethane recycling
- AU Sadadinovic, J.; Ilickovic, Z.
- CS Katedra Tehnol. Ekol., Tehnol. Fak., Sveuciliste Tuzla, Tuzla, 75000, Bosnia/Herzegovina
- SO Kemija u Industriji (2002), 51(10), 431-436 CODEN: KJUIAR: ISSN: 0022-9830
- PB Hrvatsko Drustvo Kemijskih Inzenjera i Tehnologa
- DT Journal; General Review
- LA Croatian

AB

A review. The problem of polyurethane waste recycling has major technol., ecol. and economical significance since polyurethane (PUR) is relatively expensive, and its disposal whether by burning or landfill is also costly. This paper gives the overview of existing options of PUR recycling used in industrial practice, as well as the review of alternative options of recycling of PUR waste that might be realized in practice in the near future. PUR may be recycled using phys. and chem. methods. Several phys. methods for PUR were developed such as grinding, compression molding, and adhesive pressing. Grinding has a special importance in this field, because it is the starting operation of almost all PUR recycling methods. Grinding transforms PUR foam waste into powders that can be used in the prodn. of new foams. Two methods for grinding PUR waste were developed, i.e., cryogenic grinding based on lig. nitrogen and grinding under ambient conditions. Adhesive pressing is a technique where PUR granules are mixed with PUR binder and then cured under heat and pressure. In USA, annually over 300,000 tons of elastic PUR foam scrap are recycled into carpet underlay by using this method. Some type of polyurethanes (elastomers) can be reshaped by compression molding at temps. just below the degrdn. temp. Chem. recycling based on converting of PUR polymer into starting compds. may be used to manuf. new PUR products. Most frequently used options

are glycolysis and hydrolysis. The reagent used for hydrolysis process is steam. Hydrolysis can produce polyols and amine but their reuse requires addnl. purifn. Glycolysis is a process where PUR reacts with diols (aliph. glycols) at elevated temp. to produce a glycolyzates-polyols, which can be used as a part of major polyol component in new PUR foam prodn. ICI-split phase glycolysis and BASF-method are the best known glycolysis processes used on industrial scale. Incineration with energy recovery presents a valid option of PUR recycling, esp. since the energy value of PURs is .apprx.35 MJ kg-1. Hydrogenation and pyrolysis are the promising techniques theor. applicable for PUR recycling. Those processes can break down PUR waste (and other plastics) to a valuable petrochem. feed stocks using heat, pressure, and hydrogen.

- CC 38-0 (Plastics Fabrication and Uses)
 Section cross-reference(s): 39
- ST review polyurethane urethane rubber recycling
- IT Size reduction
 - (in recycling of polyurethanes)
- IT Polyurethanes, processes
 - Urethane rubber, processes
 - (recycling of)
- IT Recycling of plastics and rubbers
 - (recycling of polyurethanes and urethane rubber)
- L58 ANSWER 3 OF 14 HCA COPYRIGHT 2008 ACS on STN
- AN 131:185680 HCA Full-text
- TI Laser pyrolysis/time-of-flight mass spectrometry studies pertinent to the behavior of flame-retarded polymers in real fire situations
- AU Price, Dennis; Gao, Fengge; Milnes, G. John; Eling, B.; Lindsay, C. I.; McGrail, P. T.
- CS Chemical Sciences Division, Science Research Institute, University of Salford, Greater Manchester, M5 $4 \rm WT$, UK
- SO Polymer Degradation and Stability (1999), 64(3), 403-410 CODEN: PDSTDW: ISSN: 0141-3910
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- AB The Salford laser pyrolysis/time-of-flight mass spectrometry
 (LP/TOFMS) technique, which models the behavior in the dark flame
 region behind the flame front in a polymer fire, was applied to study
 flame-retarded poly(Me methacrylate) (PMMA), rigid polyurethane foam
 systems, and phosphorus retarded rigid polyurethane foams and a model
 urethane compd. The laser pyrolysis of aluminum oxide trihydrate
 (ATH) retarded PMMA produces a large amt. of water and carbon dioxide
 in the volatiles. Also, the amt. of the monomer evolved is reduced

significantly compared to that obtained from pure PMMA. The implication of these results is that in a real fire situation, ATH influences PMMA pyrolysis in such a manner as to bring about a redn. in the evolved fuel while at the same time adding non-combustible gases (e.g. water) to the flame region. Thus is the PMMA flame retarded. The rigid polyurethane foams studied varied in isocyanate index and the mol. wt. of the polyols applied. The flame retardance of these materials was shown to increase with increasing isocyanate index and wt. fraction of isocyanate. Laser pyrolysis expts. of these samples showed that the major volatiles evolved were dominated by monomer and oligomers of the polypropylene glycol used to produce the foam, plus lower mol. wt. species of which carbon dioxide appeared to be a significant part. An increase in isocyanate index results in a redn. in the extent of monomer/oligomer evolution and an increase in the low mol. wt. species. With ref. to the behavior of the foams in a real fire situation, it could be imagined that the monomer/oligomer components and their breakdown products would act as fuel in the flame region while the low mol. wt. species dominated by carbon dioxide would be relatively non-flammable. An increase of isocvanate index is equiv. to making less fuel and more of the inert gases available to the burning zone and hence improving the fire resistance of the rigid polyurethane foams. The flame retardant mechanism of phosphorus, introduced as low percentages of di-Me methylphosphonate, is also attributed to a redn. in fuel evolution via pyrolysis of rigid polyurethane foams.

CC 37-5 (Plastics Manufacture and Processing)

ST polymethyl methacrylate flame retardant mechanism aluminum oxide; polywrethane foam flame retardance isocyanate index; polyol mol wt polywrethane foam flame retardance; phosphorus retarded polywrethane foam burning mechanism; laser pyrolysis mass spectrometry flame retardancy polymer

IT Combustibles

(gaseous; laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

Fire

ΙT

Fire-resistant materials

Fireproofing agents

Flammability

(laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

IT Plastic foams

Polyurethanes, processes (laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

IT Oxygen index

(limiting; laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

IT Polymer degradation

(thermal; laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

- IT 1066-53-1, Methyl methylphosphonate 21645-51-2, Aluminum oxide trihydrate, uses
 - (flame retardant; laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)

mechanism of PMMA and polyurethane foams in real fire situations)

- IT 9011-14-7, PMMA 9048-57-1, MDI-poly(propylene glycol) copolymer
 (laser pyrolysis/TOFMS studies of flame retardant
 mechanism of PMMA and polyurethane foams in
 real fire situations)
- IT 239438-61-0, n-Butyl (4 benzyl)phenylcarbamate (model urethane compd.; laser pyrolysis/TOFMS studies of flame retardant mechanism of PMMA and polyurethane foams in real fire situations)
- RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L58 ANSWER 4 OF 14 HCA COPYRIGHT 2008 ACS on STN
- AN 129:98895 HCA Full-text
- TI Polyurethane based organic aerogels and their transformation into carbon aerogels
- AU Biesmans, G.; Mertens, A.; Duffours, L.; Woignier, T.; Phalippou, J.
- CS ICI Polyurethanes, B-3078, Belg.
- SO Journal of Non-Crystalline Solids (1998), 225, 64-68 CODEN: JNCSBJ; ISSN: 0022-3093
- PB Elsevier Science B.V.
- DT Journal
- LA English
- AB New org. gels were prepd. from chem. reactions conventionally used to make polyurethane foams. Reactions were carried out using CH2C12 as solvent. Solvent exchange occurs directly in the autoclave by flushing the gel with supercrit. CO2. The subsequent org. aerogels were obtained by a classical CO2 supercrit. drying process. They are nontransparent. The thermal evolution to a carbon aerogel was

investigated with a starting polymer aerogel having a bulk d. of 0.24 g/cm3 and a sp. surface area of 300 m2/g. As the temp, increases the sp. surface area and the bulk d. increase for temps, higher than 400°C. The pore morphol. strongly depends on the temp. as evidenced by SEM expts. The pyrolyzed aerogel has the texture of an ultrafine celled foam. Thermogravimetric anal. was related to dilatometric measurements and the aerogel d. vs. temp. was estd. Carbon aerogels were obtained at temps. of 600-800°C. 57-8 (Ceramics)

Section cross-reference(s): 37

ST supercrit drying polyurethane aerogel pyrolysis transformation; polyurethane aerogel pyrolysis transformation carbon aerogel

TΤ Drving

(supercrit., carbon dioxide;

polyurethane-based org. aerogels and transformation into carbon aerogels)

RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L58 ANSWER 5 OF 14 HCA COPYRIGHT 2008 ACS on STN

AN 115:160929 HCA Full-text

OREF 115:27551a,27554a

TΙ Low-density carbonized composite foams

Kong, Fung Ming IN

PA United States Dept. of Energy, USA

U.S., 9 pp. SO CODEN: USXXAM

DT Patent

English T 70

FAN.CNT 1									
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE				
PI	US 4992254	A	19910212	US 1989-447478					
					198912 07				
				<					
	US 5047225	A	19910910	US 1990-493534					
					199003				
				<	14				
	US 5232772	A	19930803	US 1991-746528					
					199108				
					19				
				<					

PRAI US 1989-447478 A3 19891207 <--

US 1990-493534 A3 19900314 <--

The title composite foam, useful as laser targets in controlled AB fusion reactions in chromatog, columns, high-temp, catalytic supports, and structural supports, are prepd. by (a) prepg. an inverse emulsion of styrene/divinvlbenzene in water: (b) polyma, the mixt. to give a foam; (c) filling with aq. resorcinol-formaldehyde soln. inside cells of the foam; (d) curing the resorcinolformaldehyde soln. to a gel; (e) heating the resorcinol-formaldehyde gel filled polystyrene foams to a temp. sufficient to carbonize the foam. Step (c) may use a phenol-aldehyde prepolymer for the filling of the foam. Thus, the emulsion is formed from an oil phase (styrene 1.38, divinvlbenzene 1.38, and sorbitan monooleate 1.49 g) and an ag. phase (water 20.54 and sodium persulfate 0.205 g), which are mixed, heated for polymn, and crosslinking to give a foam, the pores filled consecutively with water/iso-PrOH, N2, and iso-PrOH, and then the pores filled with 2% resorcinol-formaldehyde soln. contg. Na2CO3 catalyst and water. Filling is done in vacuo, and then the foam is submerged in a resorcinol-formaldehyde soln, in a sealed bottle at 90°. Drying and pyrolysis at 1050° for 12 h in Ar, and cooling 24 h gave carbonized foam with d. 46 mg/cm3 and cell diam. <1 μ m.

IT 124-38-9, Carbon dioxide, reactions

(lig., in decontamination of low-d. carbonized composite

foams, for use in nuclear fusion)

RN 124-38-9 HCA

CN Carbon dioxide (CA INDEX NAME)

0-0-0

IC ICM C01B031-02

TNCL 423449000

CC 38-3 (Plastics Fabrication and Uses)
Section cross-reference(s): 49, 71, 80

ST carbonized crosslinked polystyrene foam; nuclear fusion target carbonized foam; building support carbonized foam; resorcinol formaldehyde phenolic resin carbonization

IT Phenolic resins, uses and miscellaneous

(crosslinked polystyrene foams contg., carbonized, for targets in nuclear fusion)

IT Carbonization and Coking

(of resorcinol-formaldehyde polymer-filled divinylbenzene-styrene copolymer foam, for target in nuclear fusion)

IT Polymerization catalysts

(sodium carbonate, for resorcinol and formaldehyde in pores of crosslinked polystyrene fram) $\,$

IΤ Building materials (support for, low d. carbonized composite foams as, prepn. of) ΙT Nuclear fusion (targets for, low d. carbonized composite feams as, prepn. of) ΙT Drying (supercrit., in prepn. of low-d. carbonized composite foams, as targets in nuclear fusion) 497-19-8, Sodium carbonate, uses and miscellaneous TT (catalyst, for polymn. of resorcinol and formaldehyde in pores of crosslinked polystyrene foam) 25014-41-9, Polyacrylonitrile 25212-86-6 ΙT (crosslinked polystyrene foams filled by, carbonized, for targets in nuclear fusion) 24969-11-7P, Resorcinol-formaldehyde resin ΤТ (formation of, in crosslinked polystyrene foam pores, for carbonized low d. composite films in nuclear fusion) ΙT 10028-17-8, Tritium, properties (lig., filling of low d. carbonized composite foams with deuterium and, in nuclear fusion) ΙT 124-38-9, Carbon dioxide, reactions (lig., in decontamination of low-d. carbonized composite foams, for use in nuclear fusion) L58 ANSWER 6 OF 14 HCA COPYRIGHT 2008 ACS on STN AN 115:94043 HCA Full-text OREF 115:16189a,16192a Low-density resorcinol-formaldehyde resin aerogels and their TT manufacture IN Pekala, Richard W. PA United States Dept. of Energy, USA SO U.S., 8 pp. Cont.-in-part of U.S. 4,873,218. CODEN: USXXAM DT Patent LA English FAN.CNT 2 PATENT NO. KIND DATE APPLICATION NO. DATE _____ ----_____ PI US 4997804 A 19910305 US 1989-406009 198909 12 <--US 406009 A0 19911115

US 199404

A0

19890901 US 1988-199404

198805

US 4873218

Α 19891010 A 2 19880526 <-- <--

PRAI US 1988-199404

AB The title materials obtained by polymn. of resorcinol with HCHO under alkali conditions give surface-functional polymer clusters which are crosslinked to give gels, which, under supercrit. conditions, form low-d. aerogels consisting of interconnected colloidal particles of diam. 100 Å. The aerogels can be carbonized to give low-d. foams with cell size 0.1 um. Addn. of a metal salt promotes the formation of a C/metal composite with catalytic properties. Thus, polymn. of resorcinol with HCHO in presence of Na2CO3 and CF3CO2H and drying in lig. CO2 for 4 h gave aerogels. Addn. of chloroplatinic acid or PdC12 in DMF to the aerogels prior to solvent exchange gave compns., which were supercrit, dried to give foams with Pt or Pd distributed throughout and then pyrolyzed at 500-1200° to give feams with high catalytic function.

B01V020-02; C08V009-28

INCL 502418000

38-3 (Plastics Fabrication and Uses) Section cross-reference(s): 37, 67

- phenolic resin manuf aerogel; resorcinol resin manuf aerogel; ST supercrit drving aerogel phenolic; carbonization phenolic resin aerogel; org foam phenolic resin aerogel; palladium contg aerogel catalyst; platinum contg aerogel catalyst
- ΙT Carbonization and Coking

(of phenolic aerogels, to low-d, carbon feams ١

- L58 ANSWER 7 OF 14 HCA COPYRIGHT 2008 ACS on STN
- AN 98:144276 HCA Full-text

OREF 98:21997a,22000a

- Study of the thermal behavior of foam polyurethanes by TΙ pyrolysis mass spectrometry
- ΑU Khmel'nitskii, R. A.; Lukashenko, I. M.; Morozova, T. P.; Sharafanov, V. T.; Khlebnikova, M. V.
- Mosk. S-kh. Akad., Moscow, USSR CS
- Izvestiva Timiryazevskoi Sel'skokhozyaistvennoi Akademii (SO 1983), (1), 186-91 CODEN: ITSAA7; ISSN: 0021-342X

- DT Journal
- LA Russian
- Pyrolysis mass-spectrometric study of cellular polyurethane compns. AB revealed that their degrdn. at .ltorsim. 800° proceeds with and without formation of monomers for compns. PPU-3F and PPU-KF, resp., but in both cases with intense liberation of CO2 gas, indicating the random character of the process. PPU-3F was prepd. from diisocyanate

DUDEG-2 (diethylene glycol-tolylene diisocyanate copolymer) [83513-02-4] and polyester P-3 (adipic acid-glycerol-sebacic acid copolymer) [29087-60-3], and modified with red P (3.2%) and C1 (0.6%). PPU-KF was made by reaction of polyisocyanate with Lapramol 292 and Phosdiol [85243-25-0] (hydroxyethylated diethylene glycol diphosphonate).

37-5 (Plastics Manufacture and Processing) CC

ST thermal degrdn cellular polyurethane; pyrolysis mass spectrometry polyurethane

Urethane polymers, reactions ΙT

> (cellular, thermal degrdn. of, pyrolysis mass-spectrometric study of)

ΙT Polymer degradation

> (thermal, of polyurethanes, cellular, pyrolysis mass-spectrometer study of)

ΙT 9073-12-5D, reaction products with P-3, chlorine- and 85243-25-0D, reaction products with phosphorus-conta. polyisocyanate and Lapramol 292 85256-22-0D, reaction products with polyisocyanate and Phosdiol (cellular, thermal degrdn. of)

L58 ANSWER 8 OF 14 HCA COPYRIGHT 2008 ACS on STN

AN 91:6169 HCA Full-text

OREF 91:1129a,1132a

TΙ Combustion of cellular urethane

Bard, S.; Clow, K. H.; Pagni, P. J. AU

CS Mech. Eng. Dep., Univ. California, Berkeley, CA, 94720, USA SO

Combustion Science and Technology (1979), 19(3-4), 141-50 CODEN: CBSTB9; ISSN: 0010-2202

Journal DT T.A

AB

English

The properties of flames supported by pyrolysis of a com. available urethane mattress foam are described. The flame shape, flame growth, species concn., solid and gas phase temps., and mass loss were measured during a 6 min period after ignition. Functional descriptions of these empirical histories of the flame radius and height, flame spread, and mass loss rates are presented. Using a simple cone-cylinder model for the flame shape, measured temps., soot vol. fractions and H2O and CO2 concns., flame emissivities (Efl) in the homogeneous nongray approxn, were calcd. Flame soot contributes up to 95% of £fl throughout the fire history. The fraction of the total combustion energy release rate emitted as radiation avs. approx. 0.18. Approx. 1/3 of this flame radiation strikes the urethane through the lower flame cylinder; the remaining 2/3 is emitted through the upper flame cone to the surroundings. Surface emission plays a significant role in detg. the net radiative flux to the pyrelysis of urethane throughout the fire history. Predictions of the unethane pyrolysis rates, based on the model for radiant heat

transfer to the mattress, are in very good agreement with the exptl. mass loss rates. From comparison between predicted and exptl. mass pyrolysis rates, convective heating of the mattress is quite negligible.

38-7 (Elastomers, Including Natural Rubber)

urethane foam mattress pyrolysis;

combustion parameter urethane foam; flame urethane foam pyrolysis

Rubber, urethane, reactions ΙT

(combustion parameters of cellular, detn. of)

ΙT Carbon black, preparation

(formation of, in combustion of urethane foam mattress)

ΤТ Flame

ST

(shape and growth, in combustion of urethane foam mattress)

TT Combustion gases

> (water and carbon dioxide, from urethane foam mattress)

- ANSWER 9 OF 14 HCA COPYRIGHT 2008 ACS on STN L58
- AN 77:1488 HCA Full-text

OREF 77:299a,302a

- Toxic products from the combustion and pyrolysis of TΙ polyurethane foams
- Napier, D. H.; Wong, T. W. AU
- CS Dep. Chem. Eng. Chem. Technol., Imp. Coll. Sci. Technol., London, UK
- SO British Polymer Journal (1972), 4(1), 45-52 CODEN: BPOJAB; ISSN: 0007-1641
- Journal DT
- LA English
- AB The thermal decompn, products of 4 polyurethane foams heated to 220-240.deg. in atms. of N2, 6% O2 in N2, and air were subjected to ir spectroscopy and chem. anal. for some of the anticipated toxic materials. When trichloroethyl phosphate [115-96-8] or 0,0-diethyl N, N-bis(2-hydroxyethyl)aminomethylphosphon ate [2781-11-5][HOCH2CH2NHCH2P(o)(OEt)2] was added to or chem. incorporated into the foams, phosphorus [7723-14-0]-contq. compds. were evolved under most of the exptl. conditions. Hydrogen cyanide [74-90-8], isocyanate, urea [57-13-6], carbon dioxide [124-38-9], halogenated compds., and alkenes were also detected. The estd. concn. of degradation products used in these tests was .sim.0.5 ppm.
- ΙT 75-13-8 124-38-9, uses and miscellaneous
- (combustion gas, from polyurethane foams)
- 75-13-8 HCA RN
- Isocvanic acid (6CI, 8CI, 9CI) (CA INDEX NAME) CN

0-0

124-38-9 HCA

Carbon dioxide (CA INDEX NAME)

RN

CN

CC 4-3 (Toxicology) ST phosphorus flame retardant toxicity; polyurethane foam thermal decompn; toxicity polyurethane foam ΤТ Alkenes, uses and miscellaneous Halogen compounds (combustion gas, from polyurethane foams) ΙT Combustion gases (from polyurethane foams, toxicity of) ΤТ 57-13-6, uses and miscellaneous 74-90-8 75-13-8 124-38-9, uses and miscellaneous (combustion gas, from polyurethane foams) ΙT 26680-22-8 (feams, combustion gases of, toxicity of) L58 ANSWER 10 OF 14 HCA COPYRIGHT 2008 ACS on STN 75:51076 HCA Full-text AN OREF 75:8075a,8078a Effect of smoke on the production and stability of high-expansion TΙ ΑU Alvares, N. J.; Lipska, A. E. CS Stanford Res. Inst., Menlo Park, CA, USA West. States Sect., Combust. Inst. [Pap.] (1971), No. SO 71-1, 31 pp. CODEN: WSCPAH DT Report. LA English AB High-expansion fire-fighting foams are produced from solns. With low surface tension and high surface viscosity. Foam formation can be inhibited by products of both combustion and pyrolysis. Foam-breaking constituents are identified from a broad class of fuels-wood, paper,

JP-5, Av-gas, and NFSO. An aq. soln. of foam conc. was sprayed onto a net and exposed to a large vol. of air flows. This produced a continuous foam mass, highly expanded and long lived. Combustion and pyrolysis products and their chem. constituents were introduced into

the air flow system, flow rates and foam degradation being controlled and measured. Pyrolysis products from all fuels tested destroy foam more effectively than combustion products, even though the latter are produced much faster. The pyrolysis products from wood are the most effective foam destroyers of all fuels tested. They are rich in chem. fractions, i.e. aldehydes, whereas combustion gives mostly CO2 and H2O. The concn. of the more effective foam breakers need be only ≤20 ppm. The fuel designations are not explained further.

50 (Propellants and Explosives) CC

fire fighting foam smoke effect ST

ΤТ Fires

(extinguishing of, smoke effects on foams for)

ΙT Foam

ΙT

(fire-extinguishing, smoke effects on)

Smoke

(foam stability in relation to, fire-extinguishing)

Gasoline, compounds IΤ

(pyrolysis products of aviation, stability of fire-extinguishhing foam in relation to)

ΙT Fuels, jet

Paper

Wood

(pyrolysis products of, stability of fire-extinguishing foam in relation to)

L58 ANSWER 11 OF 14 HCA COPYRIGHT 2008 ACS on STN

AN 72:56132 HCA Full-text

OREF 72:10301a,10304a

TΙ Rigid foam: manufacture and properties

Buist, J. M.; Hurd, R.; Stafford, R. L. AU

SO Advan. Polyurethane Technol. (1968), 187-250. Editor(s): Buist, J. M. Publisher: John Wiley and Sons Inc., New York, N. Y. CODEN: 17FIAH

Conference; General Review DT

LA

English AB A review, with 23 refs., includes the history of rigid-foam production, evaluation of resin and isocyanate starting materials, and adjunct materials, i.e. blowing agents, surfactants, catalysts, crosslinking agents, and flame-retardant additives, as well as preblending compounding problems. Flow, pressure development, and iig dwell time processing factors, and phys. properties of rigid polyurethane foams: internal pressure, thermal cond., dimensional stability, mech. properties, coeff. of expansion, esp. in building panels, ignition, burning, pyrolysis, and product fire resistance are discussed. Tests for closed-cell content, liq. H2O content, H2Owaper transmission, d., tensile strength and modulus, and shear

properties are described, and a list of rigid foam machinery manufacturers is given.

36 (Plastics Manufacture and Processing)

ST foam rigid review; review rigid foam; blowing agent foam; surfactant rigid foam; catalyst rigid foam; crosslinking agent foam; flame retardant foam; polyurethane foam rigid; thermal cond polyurethane foam; mech property polyurethane foam; fire resistance polyurethane foam; ignition polyurethane foam; water vapor transmission foam

ΙT Urethane polymers, preparation (cellular, manuf. of rigid)

1.58 ANSWER 12 OF 14 HCA COPYRIGHT 2008 ACS on STN AN 72:3988 HCA Full-text

OREF 72:751a.754a

TΙ Evolution of toxic gases from heated plastics

Bott, Barry; Firth, Jack G.; Jones, Thomas Alwyn AU

CS Safety Mines Res. Estab., Min. Power, Sheffield, UK SO British Polymer Journal (1969), 1(5), 203-4

CODEN: BPOJAB; ISSN: 0007-1641

Journal DT

LA English

The amt. and compn. of the gases evolved during the thermal decompn. AB of polyurethane (I), wrea-HCHO resin foam (II), nylon, and polyacrylonitrile (III) in air and in N was studied. Product gases were analyzed qual. by ir and mass spectral methods, and quant. by colorimetry. The main products were HCN, NH3, CO, N oxides, CO2, and H2O. The emission rate of each gas initially increased slowly with temp., but at a crit. temp. began to increase rapidly. The threshold temps, for the evaluation of each gas from each polymer were given. The wt. fraction of the polymers evolved as HCN at 500° and the activation energies for HCN and CO evolution were detd. (polymer, wt. % H CN in air, wt. % HCN in N, kJ/mole activation energy for CO in air, kJ/mole activation energy for HCN in air, kJ/mole activation energy for CO in N, and kJ/mole activation energy for HCN in N given): I, 0.57, 0.2, 67, 75, 64, 73; I, 1.35 1.4, 76, 95, 74, 73; II, 1.4, 0.2, 40, 34, 71, 42; nylon, 0.04, 0.03, 75, 56, 40, 38; III, 0.5, 0.4, 19, 42, 67, 44. All polymers evolved both HCN and NH3, indicating that C-N bonds underwent > 1 reaction in pyrolysis. The similarity of the activation energies for HCN evolution in air and N suggested that the gas evolution mechanism was not affected by oxidn. reactions. The activation energies for CO evolution in air and N were similar when the polymer mols, contained large amts, of bound O. 36 (Plastics Manufacture and Processing)

IT Activation energy
(of gas formation in pyrolysis of polymers)

Nylon, reactions
Urethane polymers, reactions
(pyrolysis of, toxic gas formation in)

IT Urea condensation products, reactions
(pyrolysis of, toxic gas formation in)

IT 74-90-8P 630-08-0P, preparation

(formation of, in pyrolysis of polymers)

IT 25014-41-9, reactions
(pyrelysis of, toxic gas formation in)

L58 ANSWER 13 OF 14 HCA COPYRIGHT 2008 ACS on STN AN 70:20597 HCA Full-text

OREF 70:3859a,3862a

 $\ensuremath{\mathsf{TI}}$ $\ensuremath{\mathsf{Bromine}}$, chlorine, and phosphorus compounds as flame retardants in rigid polyurethane foam

AU Burgess, Paul E., Jr.; Hilado, Carlos J.; Proops, William R.

CS Res. and Develop. Dep., Union Carbide Corp., South Charleston, WV, USA

- SO Space Mil. Appl. Cell. Plast. Syst., Annu. Conf., Cell. Plast. Div., Soc. Plast. Ind., 12th (1967), 3-C-1-3-C-16 Publisher: Soc. of the Plast. Ind., Inc., New York, N. Y. CODEN: 20KNAL
- DT Conference
- LA English
- The ease of ignition, flame spread, fire endurance, fuel AB contribution, and smoke of 70 polyether polyurethane rigid foam contg. tetrabromoethane, (C1CH2CH2O) 3PO, hexachlorobutadiene, or Arochlor 1254 flame retardants were used to evaluate the effect of the retardance and the usefulness of the various tests. The effect of the flame retardants on flammability performance varied with the polymer structure, flame retardant, and the flammability test used. Sucrose-based foam gave the best performance under conditions permitting pyrolysis of the carbohydrate structure to yield a carbonaceous char and water vapor to aid in flame extinguishment. Aromatic-based foam performed best when they melted prior to pyrolysis. Br-contg. retardants had excellent flame retardant properties due to the lower dissocn. energy of the C-Br bond and the high d. of the Br compds. in the volatiles evolved, but Br concns. >44.5% resulted in loss of mech. strength. Increases in the flame retardant which produced no significant effect in some flammability tests produced improvements in other tests due to the difference in the relative ability of the test to det. fire resistance under different exposure conditions.
- CC 36 (Plastics Manufacture and Processing)
- ST bromine contg flame retardants; polyurethane foam

```
(cellular, fireproofing agents for)
    Ethers
       (poly-, fire-resistant unethane polymer foam
       contq.)
    Fire-resistant materials
       (arethane polymer foam as, fireproofing
       agents for)
    87-68-3 115-96-8 11097-69-1, Aroclor 1254 25167-20-8
        (fire-resistant urethane polymer foam contg.)
TΤ
    57-50-1, Sucrose
       (reaction products with propylene oxide, fire-resistant
       urethane polymer foam contg.)
    75-56-9, Propylene oxide
       (reaction products with sucrose, fire-resistant urethane
       polymer foam contg.)
L58 ANSWER 14 OF 14 HCA COPYRIGHT 2008 ACS on STN
AN 66:85696 HCA Full-text
OREF 66:16039a,16042a
TI Porous silver catalysts
PA Erdoelchemie G.m.b.H.
SO Neth. Appl., 8 pp.
    CODEN: NAXXAN
DT Patent
T.A
    Dutch
FAN.CNT 1
    PATENT NO. KIND DATE APPLICATION NO.
                                                             DATE
    _____
                      ----
                              -----
    -----
PI NL 6605594
                              19661031 NL 1966-5594
                                                               196604
                                                               26
                                              <--
    DE 1284627
                                         DE
    FR 1489078
                                         FR
    GB 1157922
                                         GB
    GB 1157923
                                         GB
    GB 1157924
                                         GB
                              19650429 <--
PRAI DE
   Ag lactate (I) or I mixed with other metal lactates, such as Au, Pt,
AB
     Ba, Ca, and Ce, is decompd. by a heat treatment to give the title
```

; foam polyurethane; flame retardant

flame retardants; sucrose based foam

Urethane polymers, properties

(agents for, for urethane polymer foam)

Fireproofing

IΤ

ΙT

ΙT

TΤ

ΤТ

ΙT

foam; chlorine contq flame retardants; phosphorus contq

catalyst (II) useful for oxidn, of olefins, such as ethylene (III) and propylene, to epoxyalkanes, or dehydrogenation of alcs. to aldehydes. Optionally, the porosity of II may be increased by addn. of a blowing agent, such as lactic acid hydrazide (IV), to I. II can activate the reactions at temp. nearly 100° lower, as compared with conventional catalysts. For example, 500 q. H2O-contq. cryst. I was heated slowly and uniformly on a dry plate. When H2O begun to evolve from the melt I, the temp. was increased at a rate of 3°/min. to 180°, while the melt I was decompd. under gas evolution and converted to foamlike dark colored masses. The surface of the obtained masses was heated by an open flame to 230° to initiate the afterdecompn. exothermically. After cooling, the masses were crushed to give 250 g. II (sp. gr. 4). II was charged into a diphyl-heated reaction tube (1 m. length, 21 mm. inner diam.). A mixt. of 5.0 vol. % III, 6.72 vol. % CO2, and 88.28% O-N mixt. was fed (200 1./hr.) into the tube at 160° to give a gas mixt. of III 3.64, CO2 7.60, ethylene oxide (V) 0.96 vol. %, and 87.80% O-N mixt. The yield of V was 70.5%. Moreover, addn. of 50 g. IV to 500 g. I gave II (sp. gr. 1.5-2.5) which improved the yield of V to 73% in the same reaction conditions. B01J

CC 27 (Heterocyclic Compounds (One Hetero Atom)) IT 15768-18-0

IC

(pyrolysis of, porous silver catalysts by)